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(54) Title: PROCESS FOR PREPARING (-)PYRIDOBENZOXAZINE CARBOXYLIC ACID DERIVATIVES

(57) Abstract

The present invention provides a process for preparing optically active (-)pyridobenzoxazine carboxylic acid derivative and pharmaceutically acceptable salt thereof by employing a starting material of (+)ethyl 2-(4-chloro-5-fluoro-2-halo-3-nitobenzoyl)-3-[(1-hydroxypropyl-2(S)-yl)amino]acrylate. According to the present invention, optically active (-)pyridobenzoxazine carboxylic acid derivative can be manufactured from low-priced 4-chloro-5-fluorobenzoic acid derivative in a simple and economical manner.

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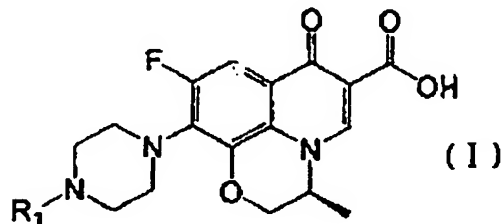
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**PROCESS FOR PREPARING (-)PYRIDOBENZOXAZINE CARBOXYLIC
ACID DERIVATIVES**

5 **BACKGROUND OF THE INVENTION**

Field of the Invention

The present invention relates to a process for preparing an optically active (-)-9-fluoro-3-methyl-7-oxo-2,3-dihydro-7H-pyrido[1,2,3-de][1,4]-benzoxazine-6-carboxylic acid derivative ("pyridobenzoxazine carboxylic acid derivative") represented by the formula(I) or pharmaceutically acceptable salt thereof having an excellent antimicrobial activity.



wherein,

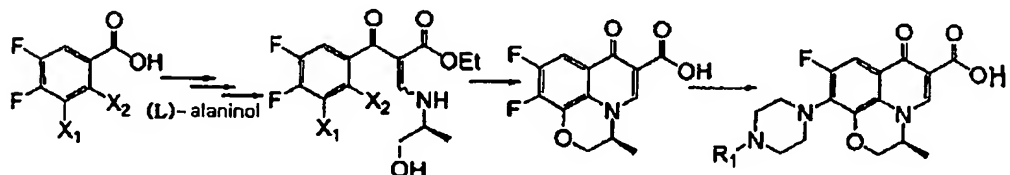
R₁ represents hydrogen atom or lower alkyl group having 1 to 5 carbon atoms.

Description of the Prior Art

A variety of optically active pyridobenzoxazine carboxylic acid derivatives have been prepared and used as active ingredients for antibiotic agents, since the compounds are known to possess higher antimicrobial activity and weaker toxicity than optically inactive racemic mixture(see: Drugs of the Future, 17(2), 559-563(1992)).

In general, optically active (-)pyridobenzoxazine carboxylic acid derivatives have been prepared in the art by the following two processes: the first one comprises a step of selective hydrolysis of (\pm)7,8-fluoro-2,3-dihydro-3-acetoxymethyl-4H-[1,4]-benzoxazine by hydrolase; and, the second one comprises a step of optical resolution of (\pm)7,8-fluoro-2,3-dihydro-3-acetoxymethyl-4H-[1,4]-benzoxazine by chemical reagent (see: EP 206,283; Korean Pat. No. 60,571).
 10 However, those processes have several drawbacks as followings: 1) theoretically 50% of isomers are lost; 2) high-priced reagent for separation is used; and, 3) complicate process of 8 steps are accompanied, which is not suitable for industrial-scale mass production. To
 15 solve the said problems, a process has been developed to prepare (-)isomer by racemizing (+)isomer obtained as a by-product during the said process (see: Japanese Patent Publication (Hei) 10-357910).

Further, processes for preparing optically active pyridobenzoxazine carboxylic acid derivatives are disclosed in U.S.Pat. Nos. 4,777,253 and 5,237,060 and Korean Pat. No. 125,115 as well. These prior arts suggest that optically active (-)pyridobenzoxazine carboxylic acid derivatives using optically active (L)-alaninol can be prepared without optical resolution, which is represented as the following reaction scheme:
 20
 25



As shown in the scheme above, a starting material of 4,5-difluorobenzoic acid derivative should be employed in the reaction, since fluorine atom among various halogen atoms is essentially required for the
 30

last step of substituting proper piperazine for 10-halogen atom. Though this process is improved in a sense that optical resolution step is not necessary, it has revealed a critical demerit that very expensive 4,5-difluorobenzoic acid derivative is required. On the other hand, it has been reported that relatively inexpensive 4-chloro-5-fluorobenzoic acid derivative, whose reactivity is lowered than 4,5-difluorobenzoic acid derivative, leads to substitution reaction at 9-fluorine atom rather than 10-fluorine atom in the last step(see: Chem. Pharm. Bull., 32, 4907-4913(1984)).

Therefore, there are strong reasons for exploring and developing a process for preparing optically active (-)pyridobenzoxazine carboxylic acid derivative by employing a low-priced material in a simple and economical manner.

SUMMARY OF THE INVENTION

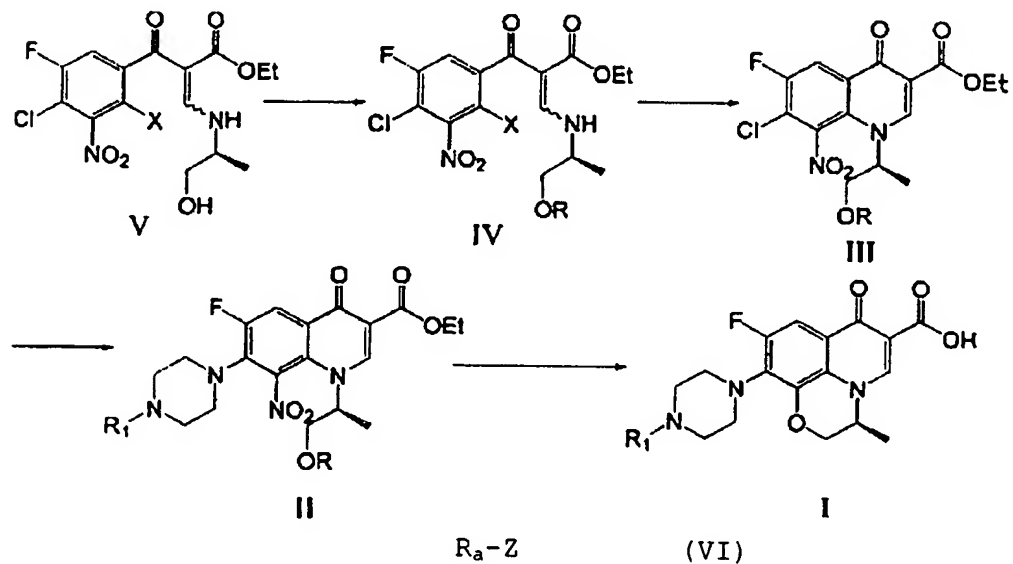
The present inventors successfully prepared optically active (-)pyridobenzoxazine carboxylic acid derivative, by employing a starting material of (+)ethyl 2-(4-chloro-5-fluoro-2-halo-3-nitobenzoyl)-3-[(1-hydroxypropyl-2(S)-yl)amino]acrylate which is obtainable from low-priced 4-chloro-5-fluorobenzoic acid derivative instead of high-priced 4,5-difluorobenzoic acid derivative, and substituting piperazine for chlorine atom.

A primary object of the present invention is, therefore, to provide a process for preparing optically active (-)pyridobenzoxazine carboxylic acid derivatives.

The other object of the invention is to provide novel compounds which are available as intermediates in the course of preparing the (-)pyridobenzoxazine carboxylic acid derivatives.

DETAILED DESCRIPTION OF THE INVENTION

In carrying out the present invention, a low-priced compound(V) is employed as a starting material which is obtainable from 4-chloro-5-fluoro-2-halo-3-nitrobenzoic acid derivatives by the known process in the art (see: U.S.Pat. No. 5,237,060). As shown in the reaction scheme below, optically active (-) pyridobenzoxazine carboxylic acid derivatives of the invention are prepared by the following steps: i) reacting a compound(V) with a reactive material(VI) or (VII) in the presence of a base to obtain a compound(IV); ii) converting the compound(IV) to a compound(III) in an organic polar solvent and in the presence of a base; iii) reacting the compound(III) with piperazine or N-monosubstituted-piperazine in an organic polar solvent in the presence of a base to obtain a novel compound(II) by so-called one-pot reaction; and, iv) hydrolyzing and cyclizing the compound(II) in an organic solvent in the presence of metal hydroxide to give the optically active compound(I).





wherein,

- X represents a halogen atom;
- Z represents a leaving group;
- Y represents an oxygen or a sulfur atom;
- 10 R_a represents -C(=O)-R₂ [wherein R₂ represents an alkyl group having 1 to 5 carbon atoms, phenyl group, substituted phenyl group, alkoxy group having 1 to 5 carbon atoms, cycloalkoxy group having 3 to 5 carbon atoms, phenoxy group, substituted phenoxy group, primary or secondary amine group or alkylthio group having 1 to 5 carbon atoms];
- 15 R_b represents alkyl group having 1 to 5 carbon atoms, phenyl group or substituted phenyl group;
- 20 R represents the same as R_a above or R_b-NH-C(=Y) [wherein R_b and Y represent the same above]; and,
- R₁ represents hydrogen atom or alkyl group having 1 to 5 carbon atoms.

25 Specifically, X includes halogen atom such as chlorine atom and fluorine atom.

Z includes halogen atom such as chloride atom and fluorine atom; carboxylate group; and, alkoxy group.

30 R₂ includes lower alkyl group having 1 to 5 carbon atoms, such as methyl group, ethyl group, n-propyl group, isopropyl group, t-butyl group, sec-butyl group, n-butyl group, isobutyl group, t-pentyl group, n-pentyl group, isopentyl group and neopentyl group, preferably methyl group and ethyl group; phenyl group; substituted phenyl group such as p-methoxyl phenyl group, 3,5-dimethoxyphenyl group, 3,5-dimethylphenyl group, 2,4,6-

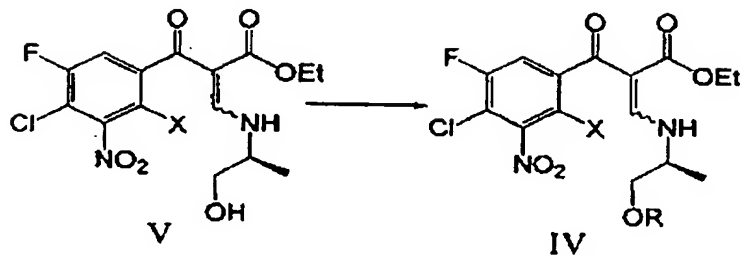
trimethylphenyl group, p-chlorophenyl group and p-fluorophenyl group; alkoxy group having 1 to 5 carbon atoms such as methoxy group, ethoxy group, n-propoxy group, t-butoxy group, sec-butoxy group, n-butoxy group,
5 isobutoxy group, t-pentoxy group, isopentoxy group, neopentoxy group and cyclopentoxy group; cycloalkoxy group having 3 to 5 carbon atoms such as cyclopropoxy group, cyclobutoxy group and cyclopentoxy group; phenoxy group; substituted phenoxy group such as p-methoxyphenoxy
10 group, p-chlorophenoxy group and p-fluorophenoxy group; primary or secondary amine group such as methylamine group, dimethylamine group, ethylamine group and diethylamine group; and, alkylthio group having 1 to 5 carbon atoms such as methylthio group, ethylthio group,
15 n-propylthiogroup, isopropylthio group, t-butylthio group, sec-butylthio group, n-butylthio group, isobutylthio group, t-pentylthio group, isopentylthio group and neopentylthio group.

R_b includes lower alkyl group having 1 to 5 carbon atoms such as methyl group, ethyl group, n-propyl group, isopropyl group, t-butyl group, sec-butyl group, n-butyl group, isobutyl group, t-pentyl group, n-pentyl group, isopentyl group and neopentyl group; phenyl group; and, substituted phenyl group such as p-methoxyphenyl group,
25 3,5-dimethoxyphenyl group, 3,5-dimethylphenyl group, 2,4,6-trimethylphenyl group, p-chlorophenyl group and p-fluorophenyl group.

R₁ includes hydrogen atom and lower alkyl group having having 1 to 5 carbon atoms such as methyl group,
30 ethyl group, n-propyl group, isopropyl group, t-butyl group, sec-butyl group, n-butyl group, isobutyl group, t-pentyl group, n-pentyl group, isopentyl group and neopentyl group.

35 The process for preparing optically active (-) pyridobenzoxazine carboxylic acid derivatives is described in more detail.

(1) Step 1: Preparation of compound(IV)



5

wherein,

X and R represent the same above.

Starting material(V) of (+)ethyl 2-(4-chloro-5-fluoro-2-halo-3-nitobenzoyl)-3-[(1-hydroxypropyl)-2(S)-yl]aminoacrylate which is obtained from 4-chloro-5-fluoro-2-halo-3-nitrobenzoic acid derivative by the conventional process (see: U.S. Pat. No. 5,237,060) is reacted with 1.0~3.0 mole equivalents of reactive material(VI) or (VII) in an organic solvent in the presence of a base at a temperature of -40°C to 80°C to obtain a compound(IV).

20 R_a-Z (VI) $R_b-N=C=Y$ (VII)

wherein,

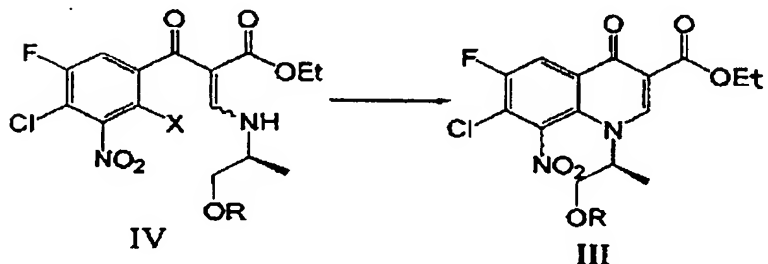
R_a, R_b, Z, and Y represent the same above.

25 The reactive material(VI) includes acylhalide, carboxylic acid anhydride, alkylchloroformate, cycloalkylchloroformate, alkylcarbonate, cycloalkylcarbonate, phenylchloroformate, substituted phenylchloroformate; and, the reactive material(VII) 30 includes isocyanate and isothiocyanate.

The base includes metal carbonate, metal bicarbonate, metal alkoxide, 1,8-diazabicyclo[5.4.0]-7-

undecene(DBU), 1,4-diazabicyclo[2.2.2]octane(DABCO), 1,5-diazabicyclo[4.3.0]-5-nonene(DBN), pyridine, dimethylaminopyridine and trimethylamine, where potassium carbonate and sodium carbonate are preferably employed as the metal carbonate; potassium bicarbonate and sodium bicarbonate, as the metal bicarbonate; and, sodium methoxide and sodium ethoxide, as the metal alkoxide.

(2) Step 2: Preparation of compound(III)



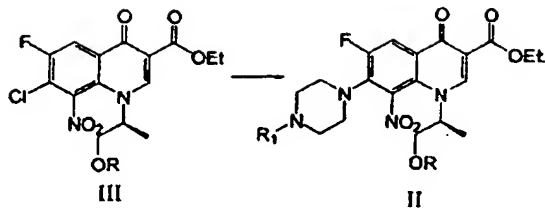
wherein,

X and R represent the same above,

15 The compound(IV) obtained in Step 1 is converted to
a compound(III) in the presence of an organic polar
solvent and 2.0~5.0 mole equivalents of base at a
temperature of range of 18°C to 150 °C depending on the
20 solvent and the base.

The organic polar solvent includes DMF(N,N' -dimethylformamide), DMSC(dimethylsulfoxide), dioxane, acetonitrile, tetrahydrofuran and acetone. The base includes metal carbonate, metal bicarbonate, metal alkoxide, DBU(1,8-diazabicyclo[5.4.0]undec-7-ene), DABCO(1,4-diazabicyclo[2.2.2]octane, DBN(1,5-diazabicyclo[4.3.0]non-5-ene), pyridine, dimethylaminopyridine and trimethylamine, where the metal carbonate, the metal bicarbonate and the metal alkoxide are the same above.

(3) Step 3: Preparation of compound (II)

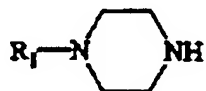


wherein,

5 X and R₁ represent the same above.

The compound(III) is reacted with 1.0~3.0 mole equivalents of piperazine or N-mono-substituted-piperazine to obtain a novel compound(II), in the presence of an organic polar solvent and 2.0~5.0 mole equivalents of a base at a temperature range of 18°C to 10 120°C. In the carrying out the said reaction, the compound(III) may be employed in a purified state or non-purified state and, the organic solvent includes DMF, 15 DMSO, dioxane, acetonitrile, tetrahydrofuran and acetone, and the base includes metal carbonate, metal bicarbonate, metal alkoxide, DBU, DABCO, DBN, pyridine, dimethylaminopyridine and trimethylamine, where the metal carbonate, the metal bicarbonate and the metal alkoxide 20 are the same above.

The piperazine or N-mono-substituted-piperazine is represented by the following formula(VIII)



25 wherein,

R₁ represents the same above.

The substituted-piperazine includes N-methylpiperazine, N-ethylpiperazine, N-n-propylpiperazine, 30 N-isopropylpiperazine, N-t-butylpiperazine, N-sec-butylpiperazine, N-n-butylpiperazine, N-

isobutylpiperazine, N-t-pentylpiperazine, N-n-pentylpiperazine, N-isopentylpiperazine and N-neopentylpiperazine.

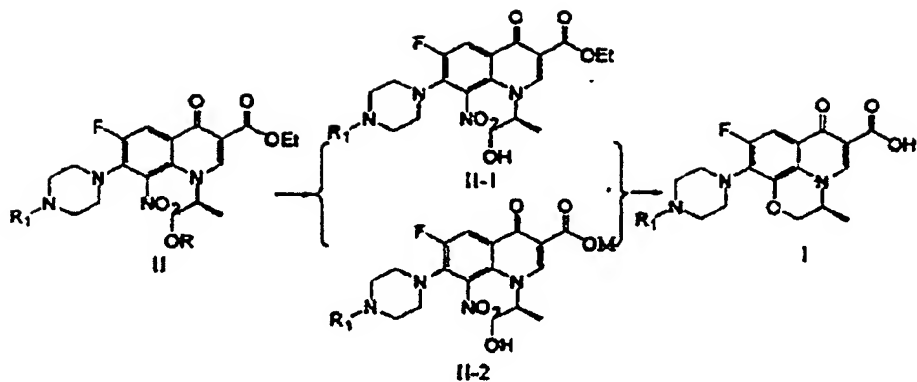
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(4) Step 4: Preparation of (-)pyridobenzoxazine carboxylic acid derivative(I)

The compound(II) is converted to a compound(I) by 10 hydrolysis and cyclization of compound(II) via one or two steps.

In carrying out the said reaction via one step, 15 the compound(I) is obtained by refluxing the compound(II) in the presence of 3.0~6.0 mole equivalents of metal hydroxide and an organic solvent with heating. The metal hydroxide includes potassium hydroxide and sodium hydroxide, and the organic solvent includes alcohol, tetrahydrofuran and a mixed solvent of one of the said solvent and water. In the case of employing the mixed 20 solvent of alcohol and water, mixing ratio may be 100:0 to 25:75(v/v), while in the case of the mixed solvent of tetrahydrofuran and water, mixing ratio of tetrahydrofuran and water, it may be 100:0 to 25:75(v/v).

In the carrying out the said reaction via two 25 steps, as shown in following reaction scheme, the compound(II) was hydrolyzed to give an intermediate compound(II-1), which is ,in turn, converted to the compound(I) by hydrolysis and cyclization of compound(II-1) in a purified or non-purified state. In addition, 30 the compound(II) was hydrolyzed to form an intermediate compound(II-2), which is ,in turn, converted to the compound(I) by cyclization of compound(II-2) in a purified or non-purified state.



wherein,

R and R₁ represent the same above; and,

M represents metal atom such as potassium and sodium.

5

The compound(II) is reacted with 1.0~2.0 mole equivalents of metal carbonate in a mixed solvent of alcohol and water to give an intermediate compound(II-1),
10 where the mixing ratio of alcohol and water in the mixed solvent may be 100:0 to 25:75(v/v), and the metal carbonate includes potassium carbonate and sodium carbonate.

Further, the compound(II) is reacted with 2.0~4.0
15 mole equivalents of metal hydroxide in alcohol to give an intermediate compound(II-2), where the metal hydroxide includes potassium hydroxide and sodium hydroxide.

The compound(I) is obtained by refluxing the
20 intermediate compound(II-1) or (II-2) in the presence of 1.0~3.0 mole equivalents of metal hydroxide and an organic solvent, where the metal hydroxide includes potassium hydroxide and sodium hydroxide, and the organic solvent includes alcohol, tetrahydrofuran and a mixed
25 solvent of one of the said solvent and water. In the case of employing the mixed solvent of alcohol and water, mixing ratio of alcohol and water may be 100:0 to 25:75(v/v), while in the case of the mixed solvent of tetrahydrofuran and water, it may be 100:0 to 25:75(v/v).

The present invention is further illustrated by the following examples, which should not be taken to limit the scope of the invention.

5

Example 1: (+) Ethyl 2-(2,4-dichloro-3-nitro-5-fluorobenzoyl)-3-[(1-acetoxyprop-2(S)-yl)amino]acrylate (IV, X=Cl, R=COMe)

10 35.0g(85mmol) of (+)ethyl 2-(2,4-dichloro-3-nitro-5-fluorobenzoyl)-3-[(1-hydroxyprop-2(S)-yl)amino]acrylate (V, X=Cl) prepared by the conventional process (see: U.S.Pat.No. 5,237,060) was dissolved in 150ml of ethylenedichloride, and chilled to a temperature
15 of -40 °C. To the resultant was added 14.3ml of triethylamine, then added 7.3ml of acetylchloride for 10 minutes at -40 °C with stirring for 1hr. Finally, to the solution was 150ml of water poured at room temperature to separate an organic layer, washed with 0.1N HCl
20 solution(50ml), 1N NaHCO₃ solution(50ml), and NaCl solution (50ml), subsequently dried over anhydrous MgSO₄, then evaporated under a reduced pressure to give 38.4g(100%, E/Z ~3/1) of the titled compound.

25 NMR(CDCl₃) δ (ppm): 10.99(q, 1H), 8.20(d, 1H), 7.17(d, 1H), 4.00-4.21(m, 5H), 2.11(s, 3H), 1.43(d, 3H), 1.04(t, 3H)

30 Example 2: (+) Ethyl 2-(2,4-dichloro-3-nitro-5-fluorobenzoyl)-3-[(1-ethoxycarboxy-propy-2(S)-yl)amino]acrylate (IV, X=Cl, R=CO₂Et)

35 17.8g(43.4mmol) of (+)ethyl 2-(2,4-dichloro-3-nitro-5-fluorobenzoyl)-3-[(1-hydroxyprop-2(S)-yl)amino]acrylate (V, X=Cl) was dissolved in 60ml of dichloroethane, and chilled to a temperature of 0° C. To the resultant was added 7.9ml of triethylamine, then a

solution obtained by dissolving 5.0ml of ethylchloroformate in 20.0ml of ethylenedichloride was added for 10 minutes at 0° C with stirring for 3hours. Finally, to the solution was 50ml of water poured at room
5 temperature to separate an organic layer, washed with 0.1N HCl solution(50ml), 1N NaHCO₃ solution(50ml), and NaCl solution (50ml), subsequently dried over anhydrous MgSO₄, then evaporated under a reduced pressure to obtain 20.93g(100%, E/Z ~3/1 of the titled compound.

10

NMR(CDCl₃) δ (ppm): 11.01(d, 1H), 8.23(d, 1H), 7.16(d,
1H), 4.00-4.29(m, 7H), 1.50(d, 3H),
1.33(t, 3H), 1.06(t, 3H)

15 Example 3: (-)Ethyl N-(1-acetoxy-propyl)-6-fluoro-
7-chloro-8-nitro-4-quinolone-3-carboxylate
(III, X=Cl, R=COMe)

20 70mg(0.15mmol) of (+)ethyl 2-(2,4-dichloro-3-nitro-
5-fluorobenzoyl)-3-[(1-acetoxypropyl)-amino]acrylate (IV, X=Cl, R=COMe) was dissolved in 2ml of acetonitrile. To the resultant was added 80mg of K₂CO₃, then refluxed for 4hours with heating. After cooling to
25 room temperature, the solvent was evaporated under a reduced pressure and treated with 5ml of acetic acid ethylester and 5ml of water to obtain organic layer, dried over anhydrous MgSO₄ and then evaporated under a reduced pressure to give 60mg(96%) of the titled compound.

30 NMR(CDCl₃) δ (ppm): 8.61(s, 1H), 8.46(d, 1H), 4.45(m, 3H),
4.31(dd, 1H), 4.13(dd, 1H), 1.94(s,
3H), 1.64(d, 3H), 1.43(t, 3H)

35 Example 4: (-)Ethyl N-(1-acetoxy-propyl)-6-fluoro-
7-(N-methylpiperazinyl)-8-nitro-4-quinolone-
3-carboxylate(II, R=COMe, R₁=Me)

60mg(0.14mmol) of (-)ethyl N-(1-acetoxy-propyl)-6-fluoro-7-chloro-8-nitro-4-quinolone-3-carboxylate(III, X=Cl, R=COMe) and 25mg of K₂CO₃ were dissolved in 3ml of acetonitrile. To the resultant was 5 added 15mg of N-methylpiperazine, then refluxed for 30 minutes with heating. After cooling to room temperature, the solvent was evaporated under a reduced pressure, then dissolved in 10ml of acetic acid ethyl ester to separate an organic layer, washed twice with 10ml of 10 water, dried over anhydrous MgSO₄, and evaporated under a reduced pressure to give 67mg(100%) of the titled compound.

NMR(CDCl₃) δ (ppm): 8.53(s, 1H), 8.31(d, 1H), 4.51(m, 1H), 15 4.39(q, 2H), 4.28(dd, 1H), 4.12(dd, 1H), 3.24(dd, 2H), 3.13(dd, 2H), 2.48(ds, 4H), 2.33(s, 3H), 1.94(s, 3H), 1.58(d, 3H), 1.40(t, 3H)

20 Example 5: (-)Ethyl N-(1-acetoxy-propyl)-6-fluoro-7-(N-methylpiperazinyl)-8-nitro-4-quinolone-3-carboxylate(II, R=COMe, R₁=Me)

54.5g(0.12mol) of (+)ethyl 2-(2,4-dichloro-3-nitro-25 5-fluorobenzoyl)-3-[(1-acetoxypropyl)-amino]acrylate (IV, X=Cl, R=COMe) was dissolved in 360ml of acetonitrile. To the resultant was added 41.6g of K₂CO₃, then refluxed for 8 hours with heating. After the starting material disappeared from TLC, 14.7ml of N-30 methylpiperazine was added to the solution slowly for 10 minutes, further refluxed for 30 minutes with heating and cooled to room temperature. Then, inorganic salt was removed by filtration and evaporated under a reduced pressure. and treated with 250ml of acetic acid ethyl 35 ester and 250ml of water to fractionate an organic layer. The organic layer was dried over anhydrous MgSO₄, evaporated under a reduced pressure to give 52g(90%) of

the titled compound. The compound was further purified by dissolving in 150ml of ethylacetate/hexane(1/2, v/v) with heating and leaving to stand at room temperature, finally to give 32g(56%) of the pure titled compound.

5

NMR(CDCl₃) δ (ppm): 8.53(s, 1H), 8.31(d, 1H), 4.51(m, 1H),
4.39(q, 2H), 4.28(dd, 1H), 4.12(dd,
1H), 3.24(dd, 2H), 3.13(dd, 2H),
2.48(ds, 4H), 2.33(s, 3H), 1.94(s,
3H), 1.58(d, 3H), 1.40(t, 3H)

10

Example 6: (-)Ethyl N-(1-ethoxycarboxy-propyl)-6-fluoro-7-(N-methylpiperazinyl)-8-nitro-4-quinolone-3-carboxylate(II, R=CO₂Et, R₁=Me)

15

20.93g(45.3mmol) of (+)ethyl 2-(2,4-dichloro-3-nitro-5-fluorobenzoyl)-3-[(1-ethoxycarboxy-propyl)-amino]acrylate(IV, X=Cl, R=CO₂Et) was dissolved in 130ml of acetonitrile. To the resultant was added 12.0g
20 of K₂CO₃, and refluxed for 8 hours with heating. Then, 5.3ml of N-methylpiperazine was further added, refluxed for 30 minutes with heating and cooled to room temperature. After the solvent was completely evaporated under a reduced pressure, dissolved in ethylacetate,
25 washed with NaCl solution, dried over anhydrous MgSO₄, and further evaporated under a reduced pressure to give 12.7g(57%) of the titled compound.

30

NMR(CDCl₃) δ (ppm): 8.56(s, 1H), 8.29(d, 1H), 4.55(m, 1H),
4.39(q, 2H), 4.36(dd, 1H), 4.23(dd,
1H), 4.11(q, 2H), 3.24(ds, 2H),
3.18(ds, 2H), 2.49(ds, 4H), 2.34(s,
3H), 1.69(d, 3H), 1.40(t, 3H),
1.21(t, 3H)

35

Example 7: (-)9-fluoro-3(S)-methyl-10-(N-methyl-1-piperazinyl)-7-oxo-2,3-dihydro-7H-pyrido

[1,2,3-de][1,4]benzoxazine-6-carboxylic acid(I, R₁=Me)

3.2g of (+)ethyl N-(1-acetoxy-propyl)-6-fluoro-7-(N-methylpiperazinyl)-8-nitro-4-quinolone-3-carboxylate(II, R=COMe, R₁=Me) was dissolved in 48ml of ethanol. To the resultant was added 2.25g of potassium hydroxide, refluxed for 2 hours with heating. Then, the solvent was evaporated under a reduced pressure, and 6.7ml of 3M AcOH solution was added to obtain a pale yellow precipitate, and added 10ml of THF while stirring. Then, the resultant solid was filtered, washed with water/THF(1/1, v/v) followed by drying to give 1.36g(57%) of the titled compound.

15

NMR(CDCl₃) δ (ppm): 14.99(s, 1H), 8.62(s, 1H), 7.74(d, 1H), 4.49(dd, 2H), 4.35(dd, 1H), 3.43(m, 4H), 2.60(d, 4H), 2.39(s, 3H), 1.63(d, 3H)

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Example 8: (-)Ethyl N-(1-hydroxy-propyl)-6-fluoro-7-N-methylpiperazinyl)-8-nitro-4-quinolone-3-carboxylate(II-1, R=H)

25 1.38g (10mmol) of K₂CO₃ was dissolved in 10ml of water, and added 2.39g(5mmol) of (-)ethyl N-(1-acetoxy-propyl)-6-fluoro-7-N-methylpiperazinyl)-8-nitro-4-quinolone-3-carboxylate(II, R=COMe, R₁=Me). 7.5ml of methanol was added and stirred for 1.5 hours at room 30 temperature. To the precipitate thus obtained was added 10ml of water, and subsequently filtered and washed with water. Then, the resultant solid was dried to give 2.1g(96%) of the titled compound.

35 NMR(CDCl₃) δ (ppm): 8.72(s, 1H), 7.74(d, 1H), 4.46(m, 1H), 4.37(q, 2H), 4.19(m, 1H), 3.92(m, 1H), 3.75(m, 2H), 3.25(ds, 2H), 3.14(ds, 2H),

2.52(ds, 4H), 2.37(s, 3H), 1.64(d, 3H),
1.40(t, 3H)

5 Example 9: (-) Potassium N-(1-hydroxy-propyl)-6-fluoro-7-N-methylpiperazinyl)-8-nitro-4-quinolone-3-carboxylate(II-2, R=H)

10 0.935g (15mmol) of KOH was dissolved in 18ml of 95% ethanol, added 2.39g(5mmol) of (-)ethyl N-(1-acetoxy-propyl)-6-fluoro-7-N-methyl-piperazinyl)-8-nitro-4-quinolone-3-carboxylate(II, R=COMe, R₁=Me), and stirred for 2 hours at room temperature. Then, the precipitate thus obtained was filtered, washed with 10ml of 95% ethanol and the resultant solid was dried to give
15 2.07g(93%) of the titled compound.

20 NMR(D₂O) δ (ppm): 8.45(s, 1H), 8.14(d, 1H), 4.28(m, 1H), 3.67(d, 2H), 3.21(ds, 2H), 3.07(ds, 2H), 3.14(ds, 2H), 2.46(ds, 4H), 2.18(s, 3H), 1.42(t, 3H)

25 Example 10: (-) 9-fluoro-3(S)-methyl-10-(N-methyl-piperazinyl)-7-oxo-2,3-dihydro-7H-pyrido[1,2,3-de][1,4]benzoxazine-6-carboxylic acid(I, R₁=Me)

30 5.1g(11.42mmol) of (-)potassium N-(1-hydroxy-propyl)-6-fluoro-7-N-methylpiperazinyl)-8-nitro-4-quinolone-3-carboxylate(II-1, R=H) was dissolved in 34ml of methanol. To the solution was added 1.07g of potassium hydroxide, then refluxed for 2.5 hours with heating. The solvent was evaporated under a reduced pressure, then 5.7ml of 3M AcOH solution was subsequently added to give a pale yellow precipitate,
35 and added 10ml of THF while stirring. Then, the resultant solid was filtered, washed with water/THF(1/1,v/v) and followed by drying to give

3.0g(73%) of the titled compound.

NMR (CDCl₃) δ (ppm): 14.99(s, 1H), 8.62(s, 1H), 7.74(d, 1H),
4.49(dd, 2H), 4.35(dd, 1H), 3.43(m,
5 4H), 2.60(d, 4H), 2.39(s, 3H),
1.63(d, 3H)

Example 11: (-)9-fluoro-3(S)-methyl-10-(N-methyl-1-piperazinyl)-7-oxo-2,3-dihydro-7H-pyrido[1,2,3-de][1,4]benzoxazine-6-carboxylic acid(I, R₁=Me)

10 5.0g of (-)ethyl N-(1-hydroxy-propyl)-6-fluoro-7-N-methylpiperazinyl)-8-nitro-4-quinolone-3-carboxylate(II-1, R=H) was reacted in the same manner as
15 in Example 10 to give about 3.0g(73%) of the titled compound.

20 NMR (CDCl₃) δ (ppm): 14.99(s, 1H), 8.62(s, 1H), 7.74(d, 1H),
4.49(dd, 2H), 4.35(dd, 1H), 3.43(m,
4H), 2.60(d, 4H), 2.39(s, 3H),
1.63(d, 3H)

25 As clearly illustrated and demonstrated above, the present invention provides a novel process for preparing optically active (-)pyridobenzoxazine carboxylic acid derivatives or pharmaceutically acceptable salt thereofs.
According to the present invention, optically active (-)
30 pyridobenzoxazine carboxylic acid derivatives can be manufactured from the low-priced 4-chloro-5-fluoro-2-halo-3-nitrobenzoic acid in a simple and economical manner.

WHAT IS CLAIMED IS:

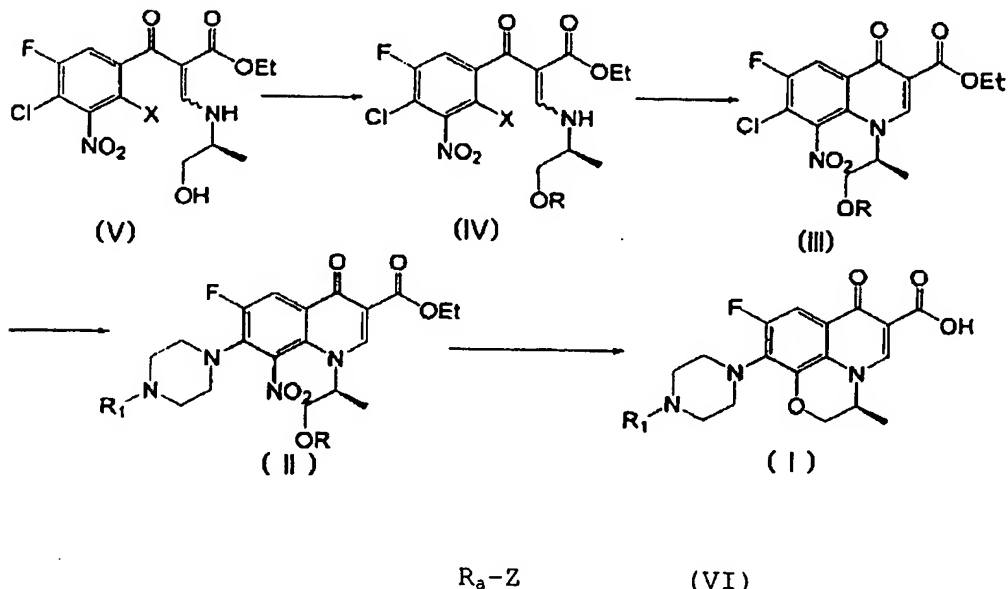
1. A process for preparing optically active (-) pyridobenzoxazine carboxylic acid derivative (I) or pharmaceutically acceptable salt thereof, which comprises the steps of:

i) reacting a compound of formula (V) with a reactive material of formula (VI) or (VII) in the presence of a base to obtain a compound of formula (IV);

10 ii) converting the compound (IV) obtained in step i) in an organic polar solvent in the presence of a base to obtain a compound of formula (III);

iii) reacting the compound (III) obtained in step ii) with piperazine or N-mono-substituted-piperazine in an organic polar solvent in the presence of a base to obtain a compound of formula (II); and,

iv) hydrolyzing and cyclizing the compound (II) obtained in said step iii) in an organic solvent in the presence of metal hydroxide to give a compound of formula (I)



25

wherein,

X represents a halogen atom;

Z represents a leaving group;

Y represents an oxygen or a sulfur atom;

5 R_a represents -C(=O)-R₂ [wherein R₂ represents an alkyl group having 1 to 5 carbon atoms, phenyl group, substituted phenyl group, alkoxy group having 1 to 5 carbon atoms, cycloalkoxy group having 3 to 5 carbon atoms, phenoxy group, substituted phenoxy group, primary or secondary amine group or alkylthio group having 1 to 5 carbon atoms];

10 R_b represents alkyl group having 1 to 5 carbon atoms, phenyl group or substituted phenyl group;

15 R represents the same as R_a above or R_b-NH-C(=Y) [wherein R_b and Y represent the same above]; and,

20 R₁ represents hydrogen atom or alkyl group having 1 to 5 carbon atoms.

2. The process of claim 1, wherein the base is selected from the group consisting of metalcarbonate, 25 metalbicarbonate, metalalkoxide, DBU(1,8-diazabicyclo[5.4.0]undec-7-ene), DABCO(1,4-diazabicyclo[2.2.2]octane, DBN(1,5-diazabicyclo[4.3.0]non-5-ene), pyridine, dimethylaminopyridine, and trimethylamine.

30 3. The process of claim 1, wherein the reactive material of formula (VI) is selected from the group consisting of acylhalide, carboxylic anhydride, alkylchloroformate, cycloalkylchloroformate, 35 alkylcarbonate, cycloalkylcarbonate, phenylchloroformate, and substituted phenylchloroformate.

4. The process of claim 1, wherein the reactive material of formula (VII) is selected from the group consisting of isocyanate and isothiocyanate

5 5. The process of claim 1, wherein the organic polar solvent in said steps ii) and iii) is selected from the group consisting of DMF(N,N'-dimethylformamide), DMSO(dimethylsulfoxide), dioxane, acetonitrile, tetrahydrofuran, and acetone.

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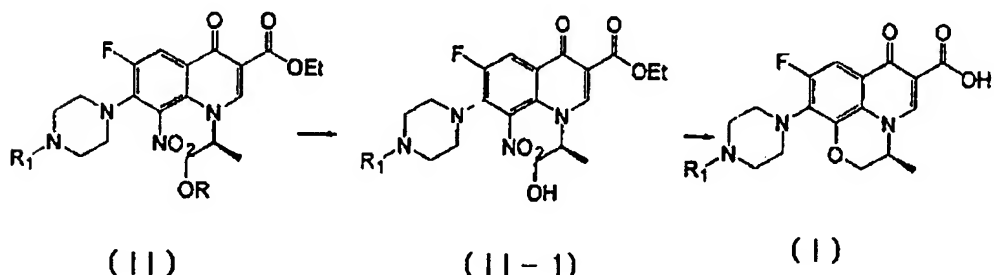
6. The process of claim 1, wherein the organic solvent in the step iv) is selected from the group consisting of alcohol, tetrahydrofuran, and a mixed solvent of one of the said solvent and water

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7. The process of claim 1, wherein the compound (III) is employed in the step iii) in a purified state or non-purified state.

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8. The process of claim 1, wherein the compound (II) is hydrolyzed to an intermediate compound of formula (II-1), which is subsequently hydrolyzed and cyclized to give the compound (I):

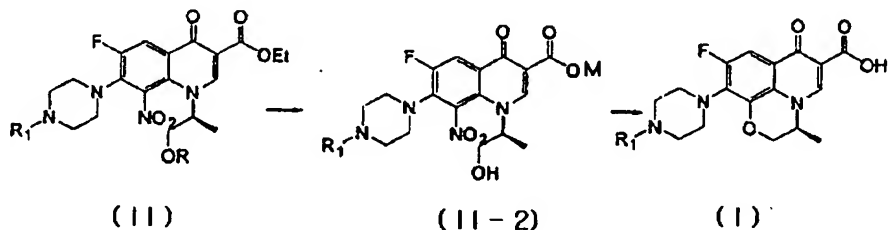


25

wherein,

R_1 represents the same above.

9. The process of claim 1, wherein the compound(II) is hydrolyzed to an intermediate compound of formula(II-2), which is subsequently cyclized to give the compound(I):



5

wherein,

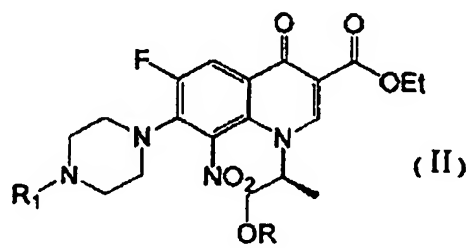
R₁ represents the same above; and,
M represents metal.

10

10. The process of claim 8, wherein the compound (II-1) is obtained by reacting the compound(II) with metal carbonate in a mixed solvent of alcohol and water.

11. The process of claim 9, wherein the compound (II-2) is obtained by reacting the compound (II) with metal hydroxide in alcohol solvent.

12. A compound of formula (II):



20

wherein,

R and R₁ represent the same above.

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13. (-) Ethyl N-(1-acetoxy-propyl)-6-fluoro-7-(N-methylpiperazinyl)-8-nitro-4-quinolone-3-carboxylate.

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INTERNATIONAL SEARCH REPORT

International application No.

PCT/KR00/00145

A. CLASSIFICATION OF SUBJECT MATTER

IPC7 C07D 498/06, C07D 265/38, C07D 241/04

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	US 477273 (ABBOTT LABORATORIES) 11 October 1988 (11. 10. 1988) see the entire document	I-13
A	JP 01-165589 A (DAIICHI SEIYAKU CO.) 29 June 1989 (29. 06. 1989) see the entire document	I-13
A	WO 90-12799 A1 (THE UPJOHN COMPANY) 1 November 1990 (01. 11. 1990) abstract; examples; claims.	I-13

 Further documents are listed in the continuation of Box C. See patent family annex.

- * Special categories of cited documents:
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- "E" earlier application or patent but published on or after the international filing date
- "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of citation or other special reason (as specified)
- "O" document referring to an oral disclosure, use, exhibition or other means
- "P" document published prior to the international filing date but later than the priority date claimed

- "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
- "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
- "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
- "&" document member of the same patent family

Date of the actual completion of the international search
31 MAY 2000 (31.05.2000)Date of mailing of the international search report
05 JUNE 2000 (05.06.2000)Name and mailing address of the ISA/KR
Korean Industrial Property Office
Government Complex-Taejon, Dunsan-dong, So-ku, Taejon
Metropolitan City 302-701, Republic of Korea
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INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No.

PCT/KR00/00145

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
US 4777253 JP 1165589	11. 10. 88 29. 06. 89	US 4826985 A JP 8026029 B4	2. 5. 89 13. 03. 96